

PCI

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau

INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5:	E	(11) International Publication Number: WO 94/03263
B01D 53/36, 53/34, 53/34 B01D 53/36, B01J 19/08 H05H 1/24, C10G 15/12	A1 (42	(43) International Publication Date: 17 February 1994 (17.02.94)
(21) International Application Number: PCT/GB93/01641	3/01641	(81) Designated States: AU, CA, JP, US, European patent (AT,
(22) International Filing Date: 4 August 1993 (04.08.93)	4.08.93)	BE, CH, DE, DK, ES, FK, GB, GK, 1E, 11, LU, MC, NL, PT, SE).
(30) Priority data: 9216509.1 4 August 1992 (04.08.92)	GB	Published With international search report. With amended claims.
(71) Applicant (for all designated States except US): PUBLIC HEALTH LABORATORY SERVICE BOARD (GB/GB); 61 Colindale Avenue, London NW9 5EQ (GB)	VBLIC D IGB/ (GB).	
(72) Inventors; and (75) Inventors; and (75) Inventors/Applicants (for US only): CLARKE, David, John (76); Intertwood, Gorandion Read, Porton, Salisbury, SP4 Oll (CB), HAYAT, Umar (GB/GB); I Sprinkfield Road, Birkby, Huddersfield HD2 2AY (GB).	id, John Salisbu- Sprink- 3).	
(74) Agent: CROPP, John, Anthony, David; Mathys & Squire, 10 Fleet Street, London EC4Y 1AY (GB).	Squire,	
(54) TItle: IMPROVEMENTS IN THE CONVERSION OF CHEMICAL MOIETIES	NOFC	HEMICAL MOIETIES

The second of th

The second secon

(57) Abstract

Association of the control of the co

ί,,

wedgested providing assess destructions as and standing of the standing of the second destruction and the second of the

Character and water

The state of the s

The second secon

A process for the conversion of a chemical moiety, which may be gaseous, liquid or a solid in fluidised form, in which the chemical moiety is reacted with a plasma or with a reagent generated by the interaction of plasma with another component, which may be a solid.



LLECTUAL PROPERTY ORGANIZATION International Bureau

ISHED UNDER THE PATENT COOPERATION TREATY (PCT)

	(11) International Publication Number:	WO 94/03263
 ¥.	A1 (43) International Publication Date:	17 February 1994 (17.02.94)

(81) Designated States: AU, CA, JP, US, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE). ·B93/01641

3 (04.08.93)

The state with the region of the state of

Published
With international search report.
With amended claims. GB

: PUBLIC ARD IGB/ Q (GB). navid, John n, Salisbu-; 1 Sprink-GB). & Squire,

ION OF CHEMICAL MOIETIES

iety, which may be gaseous, liquid or a solid in fluidised form, in which the igent generated by the interaction of plasma with another component, which

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

A Company of the

そんこくにになる ちょうに こういんしん

Las. a. 44 wermmer

nocratic People's Republic

This invention relates to improvements in or relating to processes for converting fluidised chemical moieties.

Where this activation energy is high, the reaction is to be carried out at lower pressures, e.g. atmospheric pressure or sub-atmospheric We have now found a new method of supplying the energy The reactions therefor, require less energy and are safer. The ability to carry out the reactions at lower temperatures and pressures also means that cheaper materials and simpler methods of construction can be used in the construction of In general, reactions need energy to initiate the reaction. commonly carried out at high temperature and/or pressure. and/or lower temperatures. which enables reactions the reaction vessels. pressure

with The Children or the State of the State

Alternatively improved results may be obtained at the higher temperature and/or pressure.

an improvement in processes for the conversion of a is in a fluid phase and said moiety is reacted with a In accordance with the present invention, there is provided chemical moiety characterised in that the chemical moiety plasma or with a reagent generated by the interaction of the plasma with another component. The process may involve the use of a catalyst.

is also to be understood that the invention relates to It is to be understood that the term conversion, as used herein, relates to the conversion of a material to a desired product and not merely to surface modification. It conversion of one chemical by treatment with a plasma which is not derived from the same molecule, and thus does not relate, for example, to plasma polymerisation.

	publi
	pages of pamphiets
	ö
	pages
	front
	ě
	5
}	PCT on the front

MR Mauritania MW Malawi	_			Z New Zealand		T Portugal	O Romania	_	D Sudan	E Sweden	٠,		N Schegal		-	A Ukraine	٦	_	N Vict Nam
ž	ä	ž	Ž	NZ	곱	Z	08	2	S	SE	50	SX	Š	£	2	ž	Ş	22	ž
8 £	nd Kingdom		9	r. d.	. E		-	ocratic People's Republic	жел	iblic of Korea	thsten	itenstein	ahr	mpont	•	031	Igascar		polia

Proceedable application of the communication of

ES OF INFORMATION ONLY	JNLY	
PCT on the front pages	of pamphi	PCT on the front pages of pamphlets publishing international
	2	Kanahanda
	¥	Malawi
1 Kingdom	N	Nies
	ž	Netherlands
	Q	Norway
Ž.	N N	New Zealand
	2	Poland
	ĸ.	Portugal
	9	Romania
cratic People's Republic	a	Russlan Federation
	QS	Sudan
ile of Korea	SE	Sweden
hstan	53	Slovenia
enstein	Š	Slovak Republic .
ala a	š	Senegal
pont	£	S. S.
•	2	Togo .
9	ň	Ukraine
pascar	SP	United States of America
	70	Uzbeitstan

is to improvements in or relating ting fluidised chemical moieties.

need energy to initiate the reaction.

n energy is high, the reaction is at high temperature and/or pressure.

new method of supplying the energy lons to be carried out at lower spheric pressure or sub-atmospheric wer temperatures. The ability sections at lower temperatures and that cheaper materials and simpler n can be used in the construction of

id results may be obtained at the 1/or pressure.

present invention, there is provided rocesses for the conversion of a sterised in that the chemical molety and said molety is reacted with a ent generated by the interaction of r component. The process may involve

d that the term conversion, as used the conversion of a material to a t merely to surface modification. It tood that the invention relates to ical by treatment with a plasma which he same molecule, and thus does not to plasma polymerisation.

.

WO 94/03263

7

The molety may be an element or a compound and may be gaseous or liquid or it may be a solid which is in fluidised form. Where the chemical molety is a liquid, e.g. through which the plasma is bubbled, it may be provided as such or as an aerosol, in which case the carrier gas may be or comprises the gas that is excited to plasma.

In the embodiment where the moiety is reacted with a reagent generated by the action of the plasma on another component, the plasma may be directed, for example, onto the surface of a solid to produce reactive species which react with the moiety. The solid may be a catalyst, for example. Alternatively, the reactive species may be generated from the action of the plasma on a liquid.

Plasma is normally generated from a gas; however, a liquid may also be used. For example, water may be excited to form plasmas of hydrogen and oxygen.

Any sultable means may be employed for generating the plasma. For example, it may be generated by DC glow discharge, AC electric field, plasma torch and heat, all of which may be pulsed. The heat may be generated by laser.

Alternating currents for generating the AC plasmas are preferably those having a frequency of 1-10¹³Hz, more preferably 10³-10³Hz. It will be understood, however, that in some countries the frequencies that may be used are limited, e.g. because of the risk of interference with radio transmissions. For example, in Great Britain, a frequency of 13.56MHz is set aside by the Government for such experiments and will not therefore interfere with radio transmissions. Other frequencies can be used, provided that the Government is advised of the intention to use these frequencies.

on element or a compound and may be by it may be a solid which is in the the chemical moiety is a liquid, the plasma is bubbled, it may be as an aerosol, in which case the comprises the gas that is excited to

the action of the plasma on another a may be directed, for example, onto id to produce reactive species which y. The solid may be a catalyst, for vely, the reactive species may be ition of the plasma on a liquid.

nerated from a gas; however, a liquid for example, water may be excited to igen and oxygen. may be employed for generating the e, it may be generated by DC glow c field, plasma torch and heat, all of The heat may be generated by laser.

for generating the AC plasmas are ving a frequency of 1-10¹³Hz, more It will be understood, however, that he frequencies that may be used are e of the risk of interference with For example, in Great Britain, a is set aside by the Government for will not therefore interfere with

Other frequencies can be used, srnment is advised of the intention to

WO 94/03263

c

Frequencies of less than 1 Hz may also be used. However, such frequencies may give rise to alternating or periodic glow discharge rather than a continuous plasma. Such discharges are advantageous when the power input has to be minimised or to provide additional control of the reaction.

Plasma that is generated by alternating current at radiofrequency is normally generated from gases at subatmospheric pressure. Pressures of from 100 to 10° torr are suitable. However, the pressure used is dependent on the power loadings. Therefore, if a sufficiently high power loading is available, it is possible to excite gas to plasma at a pressure above 100 Torr, if desired. However, plasmas generated by other means such as arc plasma or plasma torch are often generated at a variety of pressures ranging from sub- to super- atmospheric.

where the reaction vessel is large, as in an industrial scale reaction, it is preferable to generate the plasma at lower frequencies such as 40kHz so as to reduce the likelihood of the plasma varying in intensity across the vessel. If higher frequencies are used, nodes and antinodes of plasma intensity may be created which may result in power loss and a reduction in the efficiency of the process.

A mixture of more than one plasma may be employed and where more than one gas or liquid is excited to plasma, this may be effected before or after mixing.

While the process of the invention may be applied to conversions generally, and more particularly gaseous reactions, it is particularly useful for converting toxic gases, such as are present in internal combustion engine exhaust gases and gaseous industrial emissions, to nontoxic waste products. Either or both of the toxic gas and

WO 94/03263

Such give rise to alternating or periodic ageous when the power input has to be han 1 Hz may also be used. However, le additional control of the reaction. r than a continuous plasma.

Therefore, if a sufficiently high lable, it is possible to excite gas to ly generated from gases at sub-Pressures of from 100 to 10-3 torr r, the pressure used is dependent on ted by alternating current at radioabove 100 Torr, if desired.

h are often generated at a variety of erated by other means such as sub- to super- atmospheric.

usma varying in intensity across the and and a reduction in the efficiency of ich as 40kHz so as to reduce the essel is large, as in an industrial preferable to generate the plasma at intensity may be created which frequencies are used, nodes

one plasma may be employed and where ilquid is excited to plasma, this may after mixing.

y, and more particularly gaseous icularly useful for converting toxic resent in internal combustion engine Either or both of the toxic gas and f the invention may be applied to seous industrial emissions,

The state of the s mender in en de désidence de desidence mande de la constant

the gas employed to convert it to a non-toxic product may

be converted to plasma

from hydrocarbon burning consist mainly of CO, NOx and reaction with CO or unburned hydrocarbons to give N, and The NOx can be detoxified by Excess carbon monoxide and unburned hydrocarbon fuel Internal combustion engine exhaust gases and other exhausts are normally oxidised to CO, and water. gaseous hydrocarbons.

Section of the sectio

Examples of the detoxification of industrial gaseous emissions include the denaturing of NOx to water and nitrogen gas using hydrogen plasma, the dehalogenation of organic molecules using hydrogen plasma and the removal of fat rendering, glue and size manufacturing, tanning, fish manufacturing and cutting, food manufacturing, coffee Some of these detoxification reactions may require the odour from industrial emissions such as the emissions from polyurethane roasting, manure processing and meat processing industries. chloride and processing, polyvinyl presence of a catalyst. Exhaust gases and gaseous industrial emissions commonly process of the present invention may be used to convert the particular moieties to more acceptable gaseous products, to soluble products which can then be removed from the gas e.g. by washing, or to liquids which can be separated from the gas. For example, carbonaceous material such as soot can be treated with an oxygen plasma to form carbon include fine particulate matter dispersed in the gas.

be initiated by free radicals even in the presence of a Some reactions have such a high energy of activation that they have to be carried out at very high temperature and/or catalyst. We have now found that if such reactions are carried out in the presence of plasma in accordance with

ingine exhaust gases and other exhausts criing consist mainly of CO, NOx and s. The NOx can be detoxified by unburned hydrocarbons to give N, and ionoxide and unburned hydrocarbon fueld to CO, and water.

the denaturing of NOx to water and ydrogen plasma, the dehalogenation of lng hydrogen plasma and the removal of l emissions such as the emissions from and size manufacturing, tanning, fish olyvinyl chloride and polyurethane sutting, food manufacturing, coffee essing and meat processing industries.

ate matter dispersed in the gas. The tinvention may be used to convert the o more acceptable gaseous products, to ch can then be removed from the gas to liquids which can be separated from a, carbonaceous material such as soot than oxygen plasma to form carbon

such a high energy of activation that ed out at very high temperature and/or radicals even in the presence of a now found that if such reactions are resence of plasma in accordance with

WO 94/03263

nitro compounds, nitriles, oximes, carboxylic aromatic disproportionation Alternatively, the results achieved using such high temperature and/or free radical initiators may be improved. Reactions which may be carried out in this manner include, but are not limited to, hydrogenations such as of olefins, acetylenes, aldehydes, ketones, acids, anhydrides, esters, thereof, reductive alkylation, reductive amination, dehalogenation, migration, decomposition, carbonylation, decarbonylation, selective oxidation, acetoxylation and gas purification. this invention, the need for high temperature or compounds, anilines, phenols and derivatives ò reduced 1somerization, тау initiators hydrogenolysis,

Whilst the present invention has particular advantages when used with reactions which have previously required high temperature and/or free radical initiation, it may also be used for reactions which do not have such a high activation energy.

Many reactions are promoted by catalysts that become deactivated with the passage of time. As described in our co-pending application, entitled "Improvements in Processes Involving Catalyst", filed on the same day as the present application, such catalyst may be regenerated by contacting the surface of the catalyst with a gas in the form of a plasma. Processes may therefore be envisaged in which both the reactant mixture for a catalysed gaseous reaction and the catalyst regeneration employ plasma.

In accordance with one aspect of such processes, the catalysed reaction may take place in one time period and the regeneration of the catalyst in a second, subsequent period. Two reactors may be employed in parallel, in one of which the reaction is taking place and in the other of which the regeneration is taking place. When the catalyst in the second reactor has been regenerated, the operations

WO 94/03263

free such high carboxylic aromatic disproportionation radical initiators may be improved. ketones, acids, anhydrides, esters, carried out in this manner include, hydrogenations such as of olefins, eductive amination, dehalogenation, thereof, on, carbonylation, decarbonylation, setoxylation and gas purification. obviated. need for high temperature or phenols and derivatives achieved using Ó iles, oximes, merization, þe esults

And the second s

ntion has particular advantages when hich have previously required high ; radical initiation, it may also be h do not have such a high activation

romoted by catalysts that become ssage of time. As described in our entitled "Improvements in Processes [led on the same day as the present yst may be regenerated by contacting alyst with a gas in the form of a therefore be envisaged in which both or a catalysed gaseous reaction and on employ plasma.

take place in one time period and e catalyst in a second, subsequent hay be employed in parallel, in one is taking place and in the other of is taking place. When the catalyst as been regenerated, the operations

in the two reactors may be reversed so that the reaction is effected over regenerated catalyst in the second reactor while the catalyst of the first reactor is regenerated. Of course, more than two reactors may be used with appropriate switching arrangements.

In some cases, the gas or gases required to regenerate the from, the gaseous mixture which is to be treated in the procedure can be envisaged where in one step the gaseous mixture is treated to convert to plasma the gaseous step the same mixture is treated to convert to plasma at least one of the other gases of the mixture, being a gas The first step may also involve a reaction to generate a required gas, e.g. the gas which is to be converted to plasma, where it is not already present as such in the catalyst may already be included in, or readily generated In such cases, a self-contained employed in the regeneration of the catalyst and in another involved in the reaction which is promoted by the catalyst. gaseous components, component, or at least one of the presence of the catalyst. reaction mixture. An example of such a case is the detoxification of exhaust gas emissions from motor vehicles. For example, the catalyst employed in the catalytic converters fitted to motor vehicles for the detoxification of the exhaust gases tend to be deactivated with time due to poisoning by lead and/or phosphorus which are employed in additives for motor fuels.

Lead can be removed from the surface of the catalyst by the action of chlorine plasma which converts it to a soluble salt and phosphorus can be removed by the action of hydrogen plasma; the reactions proceeding according to the following equations:

31, -----> 2C1 (chlorine plasma)

is or gases required to regenerate the be included in, or readily generated intuctivity. In such cases, a self-contained isaged where in one step the gaseous to convert to plasma the gaseous east one of the gaseous components, eration of the catalyst and in another a is treated to convert to plasma at a gases of the mixture, being a gas ion which is promoted by the catalyst. Iso involve a reaction to generate a the gas which is to be converted to not already present as such in the

case is the detoxification of exhaust motor vehicles. For example, the the catalytic converters fitted to be detoxification of the exhaust gases and with time due to poisoning by lead chare employed in additives for motor

rom the surface of the catalyst by the lasma which converts it to a soluble can be removed by the action of reactions proceeding according to the

A CONTROL OF THE CONT

---> 2C1 (chlorine plasma)

- 7 -

WO 94/03263

Pb + 2C1 ------> PbCl₁
PbCl₂ + 2C1 -----> PbCl₄
H₂ -----> 2H (hydrogen plasma)

P + 3H ----- PH3

Thus, where halogenated compounds, for example CC1, and H₂ are present in the exhaust gases or can be generated from a gas or gases present in these gases, it will be appreciated that the regeneration of the catalyst may be achieved using the exhaust gas itself by treating it to convert one or both of the chlorine and hydrogen components thereof to plasma.

Catalytic converter systems for motor vehicles can therefore be designed wherein the catalyst is regenerated on board the vehicle, using the vehicle engine's exhaust emissions.

Where the plasma or plasmas employed for the catalyst regeneration do not interfere with the reaction which is being promoted by the catalyst, it may even be, possible to effect the catalysed reaction and the regeneration of the catalyst simultaneously.

An example of an application of the present invention to an important industrial process is in the Haber process for the catalytically promoted synthesis of ammonia from nitrogen and hydrogen. The catalyst is usually trivalent iron. Known methods require that the reaction is carried out at high temperatures and pressures such as 670% and 150 to 350 atm. Where the reaction is carried out according to the present invention, lower temperatures and pressures can be used thus reducing the risk of explosion, the energy required to carry out the synthesis and its cost.

In practice, a stoichiometric mixture of nitrogen and hydrogen is excited to plasma by any means in the presence

- 4 -

-----> PbCl₂

in the second of the second se

-> 2H (hydrogen plasma)

-----> PH3

compounds, for example CC1, and H₁ ust gases or can be generated from nt in these gases, it will be generation of the catalyst may be ust gas itself by treating it to he chlorine and hydrogen components

Control of the Contro

ystems for motor vehicles can herein the catalyst is regenerated sing the vehicle engine's exhaust lasmas employed for the catalyst erfere with the reaction which is talyst, it may even be possible to action and the regeneration of the

tion of the present invention to an oceas is in the Haber process for oted synthesis of ammonia from The catalyst is usually trivalent juire that the reaction is carried and pressures such as 670K and 150 maction is carried out according to ower temperatures and pressures can the risk of explosion, the energy the synthesis and its cost.

The second secon

ometric mixture of nitrogen and lasma by any means in the presence

WO 94/03263

1

of the catalyst to produce the ammonia. Alternatively, the admixture of hydrogen and nitrogen is excited prior to being passed over the catalyst. In this case, the catalyst will be located in a separate zone to that in which the gases are excited to plasma. The plasma is then brought into contact with the catalyst at the desired temperature and pressure. If desired, one only of the hydrogen and nitrogen is converted to plasma.

The cleavage of a carbon-carbon double bond by oxidation with ozone followed by hydrolysis to yield carbonyl compounds is an example of an application of the present invention where the moiety to be converted is a liquid. A plasma of oxygen is bubbled through a solution of the unsaturated organic compound in an inert solvent such as methanol, glacial acetic acid, ethyl acetate, hexane or chloroform at a temperature which is preferably in the region of -20°C but which may be at or above ambient temperature. The ozone is produced in the oxygen plasma.

The plasma may convert the chemical molety to a reactive substance which then takes part in a further reaction. For example, aluminium hydride may be mixed with a catalyst poisoned with sulphur and phosphorus. The mixture is exposed to a plasma of an inert gas to decompose the aluminium hydride to aluminium and hydrogen species. These species then react with the sulphur and phosphorus poisons to form a mixture of products, namely aluminium sulphide, aluminium phosphide, hydrogen sulphide and phosphine.

Alternatively, the catalyst may be mixed with zinc oxide and exposed to a hydrogen plasma. Both reactive poisons, such as mercaptan and thiol compounds, and unreactive poisons, such as aromatic sulphur compounds can be removed from the catalyst surface by this means.

 An example of the chemical molety being converted to a

by hydrolysis to yield carbonyl ple of an application of the present olety to be converted is a liquid. A bubbled through a solution of the compound in an inert solvent such as etic acid, ethyl acetate, hexane or serature which is preferably in the which may be at or above ambient ne is produced in the oxygen plasma.

it the chemical moiety to a reactive takes part in a further reaction. For ydride may be mixed with a catalyst r and phosphorus. The mixture is of an inert gas to decompose the iluminium and hydrogen species. These th the sulphur and phosphorus poisons products, namely aluminium sulphide, hydrogen sulphide and phosphine.

italyst may be mixed with zinc oxide ogen plasma. Both reactive poisons, nd thiol compounds, and unreactive atic sulphur compounds can be removed face by this means.

nemical molety being converted to a

WO 94/03263

9

reactive substance where the moiety is a liquid is where a catalyst poisoned with hydrocarbons and lead is suspended in or is in contact with dichlorine heptoxide. When the mixture is exposed to plasma of oxygen and/or inert gas, the oxygen and chlorine species formed will respectively oxidise the hydrocarbons and convert the lead to a washable lead salt.

The first of the f

h hydrocarbons and lead is suspended ere the molety is a liquid is where a

with dichlorine heptoxide. When the

plasma of oxygen and/or inert gas,

lne species formed will respectively

ons and convert the lead to a washable

WO 94/03263

phase and said moiety is reacted with a plasma or with a characterised in that the chemical moiety is in a fluid reagent generated by the interaction of plasma with another A process for the conversion of a chemical molety component.

- 2. A process according to Claim 1, wherein the moiety is in liquid form and the liquid is in the form of an aerosol.
- 3. A process according to Claim 1, wherein the chemical moiety is a fluidised finely divided solid.

- 4. A process according to Claim 5, wherein the molety is in gaseous form and is also provided in the form of plasma.
- wherein the plasma is generated by an AC electric field, by 5. A process according to any one of Claims 1 to DC glow discharge, by a laser or by plasma torch.
- 6. A process according to Claim 5, wherein the plasma is generated by an AC electric field and wherein the alternating current is supplied at from 10°Hz to 10°Hz.
- 7. A process according to Claim 5, wherein the plasma is an AC electric field and wherein the alternating current is supplied at from $10^9 \mathrm{Hz}$ to $10^{12} \mathrm{Hz}$. generated by
- A process according to any one of Claims 1 to wherein said another component is a solid.
- A process according to Claims 8, wherein said another component is a catalyst.
- 10. A process as claimed in any one of Claims 1 to 9 comprising the detoxification of a gaseous industrial

- 10 -

t the chemical moiety is in a fluid y is reacted with a plasma or with a the interaction of, plasma with another the conversion of a chemical moiety

ing to Claim 1, wherein the molety is e liquid is in the form of an aerosol.

ing to Claim 1, wherein the chemical 1 finely divided solid. ing to Claim 5, wherein the moiety is s also provided in the form of plasma. ding to any one of Claims 1 to 4, generated by an AC electric field, by g a laser or by plasma torch. ing to Claim 5, wherein the plasma is wherein the s supplied at from 103Hz to 109Hz. 3 electric field and

ing to Claim 5, wherein the plasma is 3 electric field and wherein the s supplied at from 10^{9} Hz to 10^{12} Hz.

ding to any one of Claims 1 to 7, component is a solid. ing to Claims 8, wherein said another

ment of the State 10 000 120 120 120 120 Approblem network in pipel

> laimed in any one of Claims 1 to 9 a gaseous industrial xification of

WO 94/03263

emission or internal combustion engine exhaust.

- wherein the reaction is carried out in the presence of a A process according to any one of Claims 1 to 9, catalyst. 11.
- 12. A process according to Claim II, wherein the catalyst is located in a zone remote from that in which the plasma is generated.
- wherein the reaction of the chemical molety with the plasma generates a reactive species which takes part in a second 13. A process according to any one of Claims 1 to 12, reaction.
- 14. A process according to any one of Claims 1 to 13, wherein the conversion is carried out as a continuous, semi-continuous or batch process.

ig to any one of Claims 1 to 9, carried out in the presence of a to Claim 11, wherein the catalyst ote from that in which the plasma

to any one of Claims 1 to 12, he chemical moiety with the plasma sies which takes part in a second

to any one of Claims 1 to 13, is carried out as a continuous, process.

WO 94/03263

- 12 -

PCT/GB93/01641

AMENDED CLAIMS [received by the International Bureau on 24 January 1994 (24.01.94); original claims 1-14 amended; new claims 15-25 added (3 pages)]

- A process for the conversion of a chemical moiety characterised in that the
 chemical moiety is in a fluid phase and said moiety is reacted with a plasma, or with
 a reagent generated by the interaction of plasma with another component, said
 conversion being effected in the presence of a catalyst, said catalyst being regenerated
 by in situ treatment with plasma.
- A process according to Claim 1, wherein the conversion of the chemical moiety and the catalyst regeneration are carried out simultaneously.
- A process according to Claim 1 or 2, wherein the moiety is in liquid form and the liquid is in the form of an aerosol.
- A process according to Claim 1 or 2, wherein the chemical moiety is a fluidised finely divided solid.
- 5. A process according to Claim 1 or 2, wherein the moiety is in gaseous form and is also provided in the form of plasma.
- 6. A process according to any one of Claims 1 to 5, wherein the plasma is generated by an AC electric field, by DC glow discharge, by a laser or by plasma torch.
- 7. A process according to Claim 6, wherein the plasma is generated by an AC electric field and wherein the alternating current is supplied at from 10³Hz to 10³Hz.
- 8. A process according to Claim 6, wherein the plasma is generated by an AC electric field and wherein the alternating current is supplied at from 10°Hz to 10¹²Hz.

 A process according to any one of Claims 1 to 8, wherein said another component is a solid.

PCT/GB93/01641

- 12 -

AMENDED CLAIMS ional Bureau on 24 January 1994 (24.01.94); ended; new claims 15-25 added (3 pages)]

ion of a chemical moiety characterised in that the use and said moiety is reacted with a plasma, or with teraction of plasma with another component, said presence of a catalyst, said catalyst being regenerated

n 1, wherein the conversion of the chemical moiety carried out simultaneously.

Jaim 1 or 2, wherein the moiety is in liquid form and erosol.

n 1 or 2, wherein the chemical moiety is a fluidised

n 1 or 2, wherein the moiety is in gaseous form and plasma.

ne of Claims 1 to 5, wherein the plasma is generated glow discharge, by a laser or by plasma torch.

6, wherein the plasma is generated by an AC electric current is supplied at from 10³Hz to 10³Hz. wherein the plasma is generated by an AC electric current is supplied at from 10°Hz to 10¹2Hz.

ne of Claims 1 to 8, wherein said another component

the same same color and place about

WO 94/03263

13 2

PCT/GB93/01641

10. A process according to Claim 9, wherein said another component is the catalyst.

11. A process as claimed in any one of Claims 1 to 10 comprising the detoxification of a gaseous industrial emission or internal combustion engine exhaust. 12. A process according to any one of Claims 1 to 11, wherein the catalyst is located in a zone remote from that in which the plasma is generated.

13. A process according to any one of Claims 1 to 12, wherein the reaction of the chemical moiety with the plasma generates a reactive species which takes part in a second reaction.

14. A process according to any one of Claims 1 to 13, wherein the conversion is carried out as a continuous, semi-continuous or batch process.

15. A process for the detoxification of gaseous industrial emissions or internal combustion engine exhaust characterised in that the emission or exhaust is in a fluid phase and is reacted with a plasma, or with a reagent generated by the interaction of plasma with another component, in the presence of a metallic catalyst.

16. A process according to Claim 15, wherein the emission or exhaust is in liquid form and the liquid is in the form of an aerosol.

17. A process according to Claim 15, wherein the emission or exhaust is a fluidised finely divided solid.

18. A process according to Claim 15, wherein the emission or exhaust is in gaseous form and is also provided in the form of plasma.

), wherein said another component is the catalyst.

e of Claims 1 to 10 comprising the detoxification internal combustion engine exhaust. of Claims 1 to 11, wherein the catalyst is located the plasma is generated. me of Claims 1 to 12, wherein the reaction of the enerates a reactive species which takes part in a

one of Claims 1 to 13, wherein the conversion is antinuous or batch process.

or with a reagent generated by the interaction of tion of gaseous industrial emissions or internal rised in that the emission or exhaust is in a fluid the presence of a metallic catalyst. 1 15, wherein the emission or exhaust is in liquid f an aerosol.

- 5, wherein the emission or exhaust is a fluidised
- 5, wherein the emission or exhaust is in gaseous m of plasma.

PCT/GB93/01641 . 4. WO 94/03263

19. A process according to any one of Claims 15 to 19, wherein the plasma is generated by an AC electric field, by DC glow discharge, by a laser or by plasma torch.

- 20. A process according to Claim 19, wherein the plasma is generated by an AC electric field and wherein the alternating current is supplied at from 103Hz to 103Hz.
- 21. A process according to Claim 19, wherein the plasma is generated by an AC electric field and wherein the alternating current is supplied at from 10°Hz to 1012 Hz.
- 22. A process according to any one of Claims 15 to 21, wherein said another component is a solid.
- 23. A process according to Claim 22, wherein said another component is the catalyst.
- emission or exhaust with the plasma generates a reactive species which takes part in 24. A process according to any one of Claims 15 to 23, wherein the reaction of the a second reaction.
- 25. A process according to any one of Claims 15 to 24, wherein the conversion is carried out as a continuous, semi-continuous or batch process.

Claim 19, wherein the plasma is generated by an AC alternating current is supplied at from 10³Hz to 10³Hz.

Claim 19, wherein the plasma is generated by an AC alternating current is supplied at from 10°Hz to 10¹²Hz.

, any one of Claims 15 to 21, wherein said another

laim 22, wherein said another component is the catalyst.

) any one of Claims 15 to 23, wherein the reaction of the plasma generates a reactive species which takes part in

o any one of Claims 15 to 24, wherein the conversion is semi-continuous or batch process.

INTERNATIONAL SEARCH REPORT

PCT/GB 93/01641

1,4,11 "Tr's document at particular relevance; the claimed invention cannot be contidered to introduce to considered to introduce an invention atto contract set of particular step contracts called to another set of particular step contracts to contidered as foreign as here the document is combined as foreign as here after the contract contracts and contracts to continue and the contract of the contract 1,2,8, 9,11 1,4,5, This decement published that the International filing distance or priority date and not in conflict with the application but other learnesterned the principle of theory underfulny the invention 1,2,8, 9,11-14 1,4,7, Robert to Claim No. " 08 J,C 23 C, D 53/36, "L" document member of the same patent lamily Date of Malling of this international Secret Report 01 Documentation Searched other than Minimum Documentation to the Extent that such Documents are included in the Fields Searched 4 01 J,H 05 H,C 10 G,B 29 C,C Clution of Document, 11 with indication, where appropriate, of the referent passages 44 I. CLAESIFICATION OF SUBJECT MATTER (it several classification symbols apply, Indicate all According to International Patent Charafterston (IPC) acts both National Charafterston and IPC 18 B 01 D 53/00, B 01 D 53/32, B 01 D 53/34, B 1PC 1 B 01 J 19/08, H 05 H 1/24, C 10 G 15/12 (MITSUBISHI JUKOGYO KABUSHIKI KAISHA) 09 May 1990 (09.05.90), Classification Symbols Minimum Documentation Searched ? (AMOUROUX) 17 July 1990 (17.07.90), claims. A, 5 026 949 (AMOUROUX) 25 June 1991 (25.06.91), A, 5 015 349 (SUIB) 14 May 1991 (14.05.91), T. document which may hive deadle an opticity clinical or which it clinicals in the posterior clinical another clinical enter or the resolution of a material research (a specifical).

"Or document retarting to an oral disclounts, use, stilbillian or when must "A document defining the general state of the art which is not considered to be of saricular relevance.
"E" safer socument but published on or after the international filting state. "P" document published actor to the international filling date but inter than the priestry date cistmed III. DOCUMENTS CONSIDERED TO BE RELEVANT A1, 0 366 876 Date of the Actual Compietion of the international Search A, 4 941 965 A, 3 419 597 · Special extraories of thed decuments: 10 claims. D, B claims. claims 01 ЕР, m « us, ·us, us, gs, II, FIELDS SEARCHED IV. CERTIFICATION Classification System ž, × × × × ×

Form PCTASARIO (second shoul) (January 1965)

EUROPEAN PATENT OFFICE

Signature of Authorized Office

BECKER e.h.

22. 11. 93

11 October 1993

International Application No PCT/GB 93/01641

Assurbations amobile sold, indicate sold.

12, B 01 D 53/34, B 01 D 53/36,

4, C 10 G 15/12

"unmentation Searched"

Gateficion Symbols

H,C 10 G,B 29 C,C 08 J,C 23 C,

ther than blakman Decreeration

isopropriate of the referent passages u Raiserent of Galin Na. 4

1, 2, 8,

1, 2, 8,

1, 4, 7,

91

1y 1990

GYO XABUSHIKI 10, 4, 5,

990

A service and a service to the service of

or planty data and not conflict with the septicities date or or planty data and not conflict with the specialism but closed the septicities but closed the specialism but closed the septicities and the septicities are cannot be considered to investine as the conflict and area as cannot be considered to investine as the conflict and area as cannot be considered to investine and the septicities are and area as cannot be considered to investine and the septicities are and area as cannot be considered to another and area and area and area are considered to expendite the proposition of the septicities and the septicities are more after such decreased in the art.

Date of Multing of the International Search Report
2.2. II. 93
Signature of Authorized Officer
BECKER e.h.

Company of the control of the contro

and belong and a principal of the second of

1,4,11

-2- International Application No PCT/GB 93/01641

111. BO	Chedon	111. DOCLIMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET) 111907 CKellon of Document," with backellon, where appropriate, of the referrant parasses (BEGLEY) 31 December 1968	Asterant to Claim No.	ğ	
×	DE,	(31.12.68), claims A1, 3 206 785 (VILLAMOSIPARI KUTATO INTE- ZET) 28 OCTODER 1982 (28 10, 82).	1,2		
×	ns,	the whole document. A, 4 469 508 (AMOUROUX) 04 September 1984 (04.09.84), claims.	1,3		
	·	. `		 _	
•					

Form PCT/ISA 210(extre sheet) (January 1965)

Intermettant Application No PCT/GB 93/01641

indication, where appropriate, of the relevant passages Activant to Claim No. 1,2 -13 1,3 RELEVANT (CONTINUED FROM THE SECOND SHEET) 508 } 04 September 1984 }, 785 IPARI KUTATO INTE-stober 1982 31 December 1968), document.

The best of the second of the

The same and the same of the

ANNEXE	au raport de recherche inter- national relatif à la decande de brevet international n°		La présente anneve indique les seabres de la famille de brevets cités relatifs en documents de brevets cités dans le rapport de recherche international visée ci-dessus, Les ressignements douris sont donnés à titre indicité et l'éfice.	Datus der Veröffentlichung Publication date Date de publication	22 22 22 22 22 22 22 22 22 22 22 22 22	lī	07-02-91 02-11-88 02-11-88 25-11-88 20-06-90 20-05-98 22-11-88 22-11-88 02-05-93	2001 2001 2004 2004 2004 2004 2004	rien	20-04-85 23-04-87 23-08-87 28-08-82 26-08-85 16-06-87 27-03-84 09-04-85	08-07-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-
) Patent	SAE 77571	t family tent documents ad inter- e Office is particulars the purpose	Mitglied (er) der Patent family Rember (s) Nember (s) Kembre (s) de la famile de brevets	78287 6521/89 262744 262744 26351/89 68902132 570910 370910 2639354 2639354 2639354 2639354 2639354 2639354		2861352 292391 292391 2615523 2615523 1158095 882229 172059 172059	!	- anon -	11865357 658414 3206785 184589 8201010 440046 4438706 450946	1207129 XX62499 XX62499 XX6212 S21914 S21914 S226141 S8195780 187553 187553 187553
ANNEX	to the international Search Report to the international Application No.	93/01641	This Annex lists the patent family conserver relating to the patent documents at a city and a careful to the patent documents and not not not an archive-mention. The fifter is do no may liable for these particulars and information.	Pate Pate Response	######################################	keine	NOS SERVICE OF THE PROPERTY OF	2555668 2555668	keine	SERRINGERAGE PACENTARY	40424444444444444444444444444444444444
		PCT/GB	ieder This Annex I Menbericht cited in the Ispan. national sea Iter in no way ii ibr, of inforatii	Datue der Veröffentlichung Publication date Date de publication	25-06-91	14-05-91	17-07-90		31-12-68	28-10-82	04-09-84
ANHANG	zus internationalen Rechercher- bericht über die Internationale Patentanaeldung Nr.		In diesea Anhang sind die Mitglieder IT der Phenfabalien der is bengs- nannten internationalen Recherchenbericht ti angeführten Patentdokumente angegeben, en hiese Angaben dienen nur zur Unter- richtung und erfolgen ohne Bewähr, en	la Racherchenbericht angeführtes Patentdokuaent Patent document cited in search report Document de brevet cite dans le rapport de recherche	5026949	5015349	4941965	366876	3419597	3206785	4469508
•	zwa inter bericht (Patentano		In diesea der Paten nannten i angeführt biese Ang richtung	la Rec angeführt Patent In Se Document dans le r	NS A	US A	R A	EP A1	US A	DE A1	A R
			• • •								*

Communication of the control of the

River and the editor of property and the River of